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Title:

Improvements in or Relating to Starch Content of Plants

Field of the Invention

This invention relates to novel nucleic acid sequences, vectors and host kells comprising the nucleic acid sequence(s), to polypeptides encoded thereby, and to a method of altering a host cell by introducing the nucleic acid sequence(s) of the invention.

Background to the Invention

Starch consists of two main polysaccharides, amylose and amylopectin. Amylose is a linear polymer containing α -1,4 linked glucose units, while amylopectin is a highly branched polymer consisting of a α -1.4 linked glucan backbone with α -1.6 linked glucan branches. In most plant storage reserves amylopectin consitutes about 75% of the starch content. Amylopectin is synthesized by the concerted action of soluble starch synthase and starch branching enzyme [α -1,4 glucan: α -1,4 glucan 6-glycosyltransferase, EC 2.4.1.18]. Starch branching enzyme (SBE) hydrolyses α -1.4 linkages and rejoins the cleaved glucan. via an α -1.6 linkage, to an acceptor whain to produce a branched structure. The physical properties of starch are strongly affected by the relative abundance of amylose and amylopectin, and SBE is therefore a crucial enzyme in determining both the quantity and quality of starches produced in plant systems.

Starches are commercially available from several plant sources including maize, potato and cassava. Each of these starches has unique physical characteristics and properties and a variety of possible/industrial uses. In maize there are a number of naturally occurring mutants which have altered starch composition such as high amylopectin types ("waxy" starches) or high amylose starches but in potato and cassava no such mutants exist on a commercial/basis as vet.

Genetic modification offers the possibility of obtaining new starches which may have novel and potentially useful characteristics. Most of the work to date has involved potato plants because they are amenable to genetic manipulation i.e. they can be transformed using Agrobacterium and regenerated easily from tissue culture. In addition many of the genes involved in starch biosynthesis have been cloned from potato and thus are available as targets for genetic manipulation, for example, by antisense inhibition of expression or sense suppression.

Cassava (Manihot esculenta L. Crantz) is an important crop in the tropics, where its starch-filled roots are used both as a food source and increasingly as a source of starch. Cassava is a high yielding perennial crop that can grow on poor soils and is also tolerant of drought. Cassava starch being a root-derived starch has properties similar but not identical to potato starch and is composed of 20-25% amylose and 75-80% amylopectin (Rickard et al., 1991. Trop. Sci. 31, 189-207). Some of the genes involved in starch biosynthesis have been cloned from cassava, including starch branching enzyme I (SBE I) (Salehuzzaman et al., 1994 Plant Science 98, 53-62), and granule bound starch synthase I (GBSS I) (Salehuzzaman et al., 1993 Plant Molecular Biology 23, 947-962) and some work has been done on their expression patterns although only in in vitro grown plants (Salehuzzaman et al., 1994 Plant Science 98, 53-62).

In most plants studied to date e.g. maize (Boyer & Preiss, 1978 Biochem. Biophys. Res. Comm. 80, 169-175), rice (Smyth. 1988 Plant Sci. 57, 1-8) and pea (Smith, Planta 175, 270-279). two forms of SBE have been identified, each encoded by a separate gene. A recent review by Burton et al.. (1995 The Plant Journal 7, 3-15) has demonstrated that the two forms of SBE constitute distinct classes of the enzyme such that, in general, enzymes of the same class from different plants may exhibit greater similarity than enzymes of different classes from the same plant. In their review, Burton et al. termed the two respective enzyme families class "A" and class "B", and the reader is referred thereto (and to the references cited therein) for a detailed discussion of the distinctions between the two classes. One general distinction of note would appear to be the presence, in class A SBE molecules, of a flexible N-terminal domain, which is not found in class B molecules. The distinctions noted by Burton et al. are relied on herein to define class A and class B SBE

molecules, which terms are to be interpreted accordingly.

Many organisations have interests in obtaining modified Cassava starches by means of genetic modification. This is impossible to achieve however, unless the plant is amenable to transformation and regeneration, and the starch biosynthesis genes which are to be targeted for modification must be cloned. The production of transgenic cassava plants has only recently been demonstrated (Taylor et al., 1996 Nature Biotechnology 14, 726-730; Schöpke et al., 1996 Nature Biotechnology 14, 731-735; and Li et al., 1996 Nature Biotechnology 14, 736-740). The present invention concerns the identification, cloning and sequencing of a starch biosynthetic gene from Cassava, suitable as a target for genetic manipulation.

Summary of the Invention

In a first aspect the invention provides a nucleic acid sequence encoding a polypeptide having starch branching enzyme (SBE) activity, the polypeptide comprising an effective portion of the amino acid sequences shown in Figure 4 or Figure 13. The nucleic acid is conveniently in substantial isolation, especially in isolation from other naturally associated nucleic acid sequences.

An "effective portion" of the amino acid sequences may be defined as a portion which retains sufficient SBE activity when expressed in *E. coli* KV832 to complement the branching enzyme mutation therein. The amino acid sequences shown in Figures 4 and 13 include the N terminal transit peptide, which comprises about the first 50 amino acid residues. As those skilled in the art will be well aware, such a transit peptide is not essential for SBE activity. Thus the mature polypeptide, lacking a transit peptide, may be considered as one example of an effective portion of the amino acid sequence shown in Figure 4 or Figure 13.

Other effective portions may be obtained by effecting minor deletions in the amino acid sequence, whilst substantially preserving SBE activity. Comparison with known class A SBE sequences, with the benefit of the disclosure herein, will enable those skilled in the

art to identify regions of the polypeptide which are less well conserved and so amenable to minor deletion, or amino acid substitution (particularly, conservative amino acid substitution) whilst substantially preserving SBE activity. Such less well-conserved regions are generally found in the N terminal amino acid residues (up to the triple proline "elbow" at residues 138-140 in Figure 4 and up to the proline elbow at residues 143-145 in Figure 13) and in the last 50 residues or so of the C terminal, and in particular in the acidic tail of the C terminal.

Conveniently the nucleic acid sequence is obtainable from cassava, preferably obtained therefrom, and typically encodes a polypeptide obtainable from cassava. In a particular embodiment, the encoded polypeptide may have the amino acid sequence NSKH at about position 697 (in relation to Figure 4), which sequence appears peculiar to an isoform of the SBE class A enzyme of cassava, other class A SBE enzymes having the conserved sequence DA D/E Y (Burton *et al.*, 1995 cited above).

In a particular aspect of the invention there is provided a nucleic acid comprising a portion of nucleotides 21 to 2531 of the nucleic acid sequence shown in Figure 4, or a functionally equivalent nucleic acid sequence. Such functionally equivalent nucleic acid sequences include, but are not limited to, those sequences which encode substantially the same amino acid sequence but which differ in nucleotide sequence from that shown in Figure 4 by virtue of the degeneracy of the genetic code. For example, a nucleic acid sequence may be altered (e.g. "codon optimised") for expression in a host other than cassava, such that the nucleotide sequence differs substantially whilst the amino acid sequence of the encoded polypeptide is unchanged. Other functionally equivalent nucleic acid sequences are those which will hybridise under stringent hybridisation conditions (e.g. as described by Sambrook et al., Molecular Cloning. A Laboratory Manual, CSH, i.e. washing with 0.1xSSC, 0.5% SDS at 68°C) with the sequence shown in Figure 4. Figure 10 shows a functionally equivalent sequence designated "125 + 94", which includes a region corresponding to the 3' coding portion of the sequence in Figure 4. Figure 13 shows a functionally equivalent sequence which comprises a second complete SBE coding sequence (the SBE-derived sequence is from nucleotides 35 to 2760, of which the coding sequence is nucleotides 131-2677, the rest of the sequence in the figure is vector-derived).

Functionally equivalent DNA sequences will preferably comprise at least 200-300bp, more preferably 300-600bp, and will exhibit at least 88% identity (more preferably at least 90%, and most preferably at least 95% identity) with the corresponding region of the DNA sequence shown in figures 4 or 10. Those skilled in the art will readily be able to conduct a sequence alignment between the putative functionally equivalent sequence and those detailed in Figures 4 or 10 - the identity of the two sequences is to be compared in those regions which are aligned by standard computer software, which aligns corresponding regions of the sequences.

In particular embodiments the nucleic acid sequence may alternatively comprise a 5' and/or a 3' untranslated region ("UTR"), examples of which are shown in Figures 2 and 4. Figure 9 includes a 3' UTR, as nucleotides 688-1044 and Figure 10 includes 3' UTR as nucleotides 1507-1900 (which nucleotides correspond to the first base after the "stop" codon to the base immediately preceding the poly (A) tail). Any one of the sequences defined above, or a functional equivalent thereof (as defined by hybridisation properties, as set out in the preceding paragraph), could be useful in sense or anti-sense inhibition of corresponding genes, as will be apparent to those skilled in the art. It will also be apparent to those skilled in the art that such regions may be modified so as to optimise expression in a particular type of host cell and that the 5' and/or 3' UTRs could be used in isolation, or in combination with a coding portion of the sequence of the invention. Similarly, a coding portion could be used without a 5' or a 3' UTR if desired.

In a further aspect, the invention provides a replicable nucleic acid construct comprising any one of the nucleic acid sequences defined above. The construct will typically comprise a selectable marker and may allow for expression of the nucleic acid sequence of the invention. Conveniently the vector will comprise a promoter (especially a promoter sequence operable in a plant and/or a promoter operable in a bacterial cell) and one or more regulatory signals known to those skilled in the art.

In another aspect the invention provides a polypeptide having SBE activity, the polypeptide comprising an effective portion of the amino acid sequence shown in Figure 4 or Figure 13. The polypeptide is conveniently one obtainable from cassava, although it may be

derived using recombinant DNA techniques. The polypeptide is preferably in substantial isolation from other polypeptides of plant origin, and more preferably in substantial isolation from any other polypeptides. The polypeptide may have amino acid residues NSKH at about position 697 (in the sequence shown in Figure 4), instead of the sequence DA D/E Y found in other SBE class A polypeptides. The polypeptide may be used in a method of modifying starch *in vitro*, the method comprising treating starch under suitable conditions (of temperature, pH etc.) with an effective amount of the polypeptide.

Those skilled in the art will appreciate that the disclosure of the present specification can be utilised in a number of ways. In particular, the characteristics of a host cell may be altered by recombinant DNA techniques. Thus, in a further aspect, there is provided a method by which a host cell may be altered by introduction of a nucleic acid sequence comprising at least 200bp and exhibiting at least 88% sequence identity (more preferably at least 90%, and most preferably at least 95% identity) with the corresponding region of the DNA sequence shown in Figures 4, 9, 10 or 13, operably linked in the sense or (preferably) in the anti-sense orientation to a suitable promoter active in the host cell, and causing transcription of the introduced nucleic acid sequence, said transcript and/or the translation product thereof being sufficient to interfere with the expression of a homologous gene naturally present in said host cell, which homologous gene encodes a polypeptide having SBE activity. The altered host cell is typically a plant cell, such as a cell of a cassava, banana, potato, sweet potato, tomato, pea, wheat, barley, oat, maize, or rice plant.

Desirably the method further comprises the introduction of one or more nucleic acid sequences which are effective in interfering with the expression of other homologous gene or genes naturally present in the host cell. Such other genes whose expression is inhibited may be involved in starch biosynthesis (e.g. an SBE I gene), or may be unrelated to SBE II.

Those skilled in the art will be aware that both anti-sense inhibition, and "sense suppression" of expression of genes, especially plant genes, has been demonstrated (e.g. Matzke & Matzke 1995 Plant Physiol. 107, 679-685).

It is believed that antisense methods are mainly operable by the production of antisense mRNA which hybridises to the sense mRNA, preventing its translation into functional polypeptide, possibly by causing the hybrid RNA to be degraded (e.g. Sheehy et al., 1988 PNAS 85, 8805-8809; Van der Krol et al., Mol. Gen. Genet. 220, 204-212). Sense suppression also requires homology between the introduced sequence and the target gene, but the exact mechanism is unclear. It is apparent however that, in relation to both antisense and sense suppression, neither a full length nucleotide sequence, nor a "native" sequence is essential. Preferably the nucleic acid sequence used in the method will comprise at least 200-300bp, more preferably at least 300-600bp, of the full length sequence, but by simple trial and error other fragments (smaller or larger) may be found which are functional in altering the characteristics of the plant. It is also known that untranslated portions of sequence can suffice to inhibit expression of the homologous gene - coding portions may be present within the introduced sequence, but they do not appear to be essential under all circumstances.

The inventors have discovered that there are at least two class A SBE genes in cassava. A fragment of a second gene has been isolated, which fragment directs the expression of the C terminal 481 amino acids of cassava class A SBE (see Figure 10) and comprises a 3' untranslated region. Subsequently, a complete clone of the second gene was also recovered (see Figure 12). The coding portions of the two genes show some slight differences, and the second SBE gene may be considered as functionally equivalent to the corresponding portion of the nucleotide sequence shown in Figure 4. However, the 3' untranslated regions of the two genes show marked differences. Thus the method of altering a host cell may comprise the use of a sufficient portion of either gene so as to inhibit the expression of the naturally occurring homologous gene. Conveniently, a portion of nucleotide sequence is employed which is conserved between both genes. Alternatively, sufficient portions of both genes may be employed, typically using a single construct to direct the transcription of both introduced sequences.

In addition, as explained above, it may be desired to cause inhibition of expression of the class B SBE (i.e. SBE I) in the same host cell. A number of class B SBE gene sequences are known, including portions of the cassava class B SBE (Salehuzzaman et al., 1994)

Plant Science 98, 53-62) and any one of these may prove suitable. Preferably the sequence used is that which derives from the host cell sought to be altered (e.g. when altering the characteristics of a cassava plant cell, it is generally preferred to use sense or anti-sense sequences corresponding exactly to at least portions of the cassava gene whose expression is sought to be inhibited).

In a further aspect the invention provides an altered host cell, into which has been introduced a nucleic acid sequence comprising at least 200bp and exhibiting at least 88% sequence identity (more preferably at least 90%, and most preferably at least 95% identity) with the corresponding region of the DNA sequence shown in Figures 4, 9, 10 or 13, operably linked in the sense or anti-sense orientation to a suitable promoter, said host cell comprising a natural gene sharing sequence homology with the introduced sequence.

The host cell may be a micro-organism (such as a bacterial, fungal or yeast cell) or a plant cell. Conveniently the host cell altered by the method is a cell of a cassava plant, or another plant with starch storage reserves, such as banana, potato, sweet potato, tomato, pea, wheat, barley, oat, maize, or rice plant. Typically the sequence will be introduced in a nucleic acid construct, by way of transformation, transduction, micro-injection or other method known to those skilled in the art. The invention also provides for a plant into which has been introduced a nucleic acid sequence of the invention, or the progeny of such a plant.

The altered plant cell will preferably be grown into an altered plant, using techniques of plant growth and cultivation well-known to those skilled in the art of re-generating plantlets from plant cells.

The invention also provides a method of obtaining starch from an altered plant, the plant being obtained by the method defined above. Starch may be extracted from the plant by any of the known techniques (e.g. milling). The invention further provides starch obtainable from a plant altered by the method defined above, the starch having altered properties compared to starch extracted from an equivalent but unaltered plant. Conveniently the altered starch is obtained from an altered plant selected from the group

consisting of cassava, potato, pea, tomato, maize, wheat, barley, oat, sweet potato and rice. Typically the altered starch will have increased amylose content.

The invention will now be further described by way of illustrative examples and with reference to the accompanying drawings, in which:-

Figure 1 is a schematic illustration of the cloning strategy for cassava SBE II. The top line represents the size of a full length clone with distances in kilobases (kb) and arrows representing oligonucleotides (rightward pointing arrows are sense strand, leftward are on opposite strand). The long thick arrow is the open reading frame with start and stop codons shown. Below this are shown the 3' RACE, 5' RACE and PCR clones identified either by the plasmid name (shown in brackets above the line) or the clone number (shown to the left of the clone) for the 5' RACE only. Also shown (by an x) in the 5' RACE clones are positions of small deletions or introns.

Figure 2 shows the DNA sequence and predicted ORF of csbe2con.seq. This sequence is a consensus of 3' RACE pSJ94 and 5' RACE clones 27/9,11 and 28. The first 64 base pairs are derived from the RoRidT17 adaptor primer/dT tail followed by the SBE sequence. The one long open reading frame is shown in one letter code below the double strand DNA sequence. Also shown is the upstream ORF (MQL...LPW).

Figure 3 shows an alignment of the 5' region of cassava SBE II csbe2con and pSJ99 (clones 20 and 35) DNA sequences. Differences from the consensus sequence are shaded.

Figure 4 shows the DNA sequence and predicted ORF of full length cassava SBE II tuber cDNA in pSJ107. The sequence shown is from the CSBE214 to the CSBE218 oligonucleotide. The DNA sequence is sequence ID No. 28 in the attached sequence listing; the amino acid sequence is Seq ID No. 29.

Figure 5 shows an alignment of 3' region of cassava SBE II pSJ116 and 125+94 DNA sequences. The top line is the 125 + 94 sequence and the bottom SJ116 sequence. Identical nucleotides are indicated by the same letter in the middle line, differences are

indicated by a gap, and dashed lines indicate gaps introduced to optimise alignment.

Figure 6 shows an alignment of carboxy terminal region of pSJ116 and 125+94 protein sequences. The top sequence is from 125+94 and the bottom from pSJ116. Identical amino acid residues are shown with the same letter, conserved changes with a colon and neutral changes with a period.

Figure 7 shows a phylogenetic tree of starch branching enzyme proteins. The length of each pair of branches represents the distance between sequence pairs. The scale beneath the tree measures the distance between sequences (units indicate the number of substitution events). Dotted lines indicate a negative branch length because of averaging the tree. Zmcon12.pro is maize SBE II, psstb1.pro is pea SBE I (Bhattacharyya et al 1990 Cell 60, 115-121) and atsbe2-1 & 2-2.pro are two SBE II proteins from Arabidopsis thalania (Fisher et al 1996 Plant Mol. Biol. 30, 97-108). SJ107.pro is representative of a cassava SBE II sequence, and potsbe2.pro is a potato SBE II sequence known to the inventors.

Figure 8 is an alignment of SBE II proteins. Protein sequences are indicated in one letter code. The top line represents the consensus sequence, below which is shown the consensus ruler and the individual SBE II sequences. Residues matching the consensus are shaded. Dashes represent gaps introduced to optimise alignment. Sequence identities are shown at the right of the figure and are as Figure 7, except that SJ107 pro is cassava SBE II.

Figure 9 shows the DNA sequence and predicted ORF of a cassava SBE II cDNA isolated by 3' RACE (plasmid pSJ 101).

Figure 10 shows the consensus DNA sequence and predicted ORF of a second cassava SBE II cDNA isolated by 3' and 5' RACE (sequence designated 125+94 is from plasmid pSJ125 and pSJ94, spliced at the CSBE217 oligo sequence).

Figure 11 is a schematic diagram of the plant transformation vector pSJ64. The black line represents the DNA sequence. The hashed line represents the bacterial plasmid backbone

(containing the origin of replication and bacterial selection marker) and is not shown in full. The filled triangles represent the T-DNA borders (RB = right border, LB = left border). Relevant restriction enzyme sites are shown above the black line with the approximate distances (in kiloobases) betwen sites marked by an asterisk shown underneath. The thinnest arrows represent polyadenylation signals (pAnos = nopaline synthase, pAg7 = Agrobacterium gene 7), the intermediate arrows represent protein coding regions (SBE II = cassava SBE II, HYG = hygromycin resistance gene) and the thick arrows represent promoter regions (P-2x35S = double CaMV 35S promoter, P-nos = nopaline synthase promoter).

Figure 12 is a schematic illustration of the cloning strategy used to isolate a second cassava SBE II gene. The top line represents the size of a full length clone with distances in kilobases (kb) and arrows representing oligonucleotides (rightward pointing arrows are sense strand, leftward are on opposite strand). The long thick arrow is the open reading frame with start and stop codons shown. Below this are shown the 3' RACE, 5' RACE and PCR clones identified either by the plasmid name (shown in brackets above the line) or the clone number (shown to the right of the clone).

Figure 13 shows the DNA sequence and predicted ORF of a second full length cassava SBE II tuber cDNA in pSJ146. Nucleotides 35-2760 are SBE II sequence and the remainder are from the pT7Blue vector. The DNA sequence of Figure 13 is Seq ID No. 30, and the amino acid sequence is Seq ID No. 31, in the attached sequence listing.

Example 1

This example relates to the isolation and cloning of SBE II sequences from cassava.

Recombinant DNA manipulations

Standard procedures were performed essentially according to Sambrook *et al.* (1989 Molecular cloning A laboratory manual, 2nd edn. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.). DNA sequencing was performed on an ABI automated DNA sequencer and sequences manipulated using DNASTAR software for the Macintosh.

Rapid Amplification of cDNA ends (RACE) and PCR conditions

5' and 3' RACE were performed essentially according to Frohman et al., (1988 Proc. Natl. Acad. Sci. USA 85, 8998-9002) but with the following modifications.

For 3' RACE, 5 μ g of total RNA was reverse transcribed using 5 pmol of the RACE adaptor RoRidT17 as primer and Stratascript RNAse H- reverse transcriptase (50 U) in a 50 μ l reaction according to the manufacturer's instructions (Stratagene). The reaction was incubated for 1 hour at 37°C and then diluted to 200 μ l with TE (10 mM Tris HCl, 1 mM EDTA) pH 8 and stored at 4°C. 2.5 μ l of this cDNA was used in a 25 μ l PCR reaction with 12.5 pmol of SBE A and Ro primers for 30 cycles of 94°C 45 sec, 50°C 25 sec, 72°C 1 min 30 sec. A second round of PCR (25 cycles) was performed using 1 μ l of this reaction as template in a 50 μ l reaction under the same conditions. Amplified products were separated by agarose gel electrophoresis and cloned into the pT7Blue vector (Invitrogen).

For the first round of 5' RACE, 5 μ g of total leaf RNA was reverse transcribed as described above using 10 pmol of the SBE II gene specific primer CSBE22. This primer was removed from the reaction by diluting to 500 μ l with TE and centrifuging twice through a centricon 100 microconcentrator. The concentrated cDNA was then dA-tailed with 9U of terminal deoxynucleotide transferase and 50 μ M dATP in a 20 μ l reaction in buffer supplied by the manufacturer (BRL). The reaction was incubated for 10 min at 37°C and 5 min at 65°C and then diluted to 200 μ l with TE pH 8. PCR was performed in a 50 μ l volume using 5 μ l of tailed cDNA, 2.5 pmol of RoRidT17 and 25 pmol of Ro and CSBE24 primers for 30 cycles of 94°C 45 sec, 55°C 25 sec, 72°C 3 min. Amplified products were separated on a 1% TAE agarose gel, cut out, 200 μ l of TE was added and melted at 99°C for 10 min. Five μ l of this was re-amplified in a 50 μ l volume using CSBE25 and Ri as primers and 25 cycles of 94°C 45 sec, 55°C 25 sec, 72°C 1 min 30 sec. Amplified fragments were separated on a 1% TAE agarose gel, purified on DEAE paper and cloned into pT7Blue.

The second round of 5' RACE was performed using CSBE28 and 29 primers in the first and second round PCR reactions respectively using a new A-tailed cDNA library primed

with CSBE27.

A third round of 5' RACE was performed on the same CSBE27 primed cDNA.

Repeat 3' RACE and PCR Cloning

The 3' RACE library (RoRidT17 primed leaf RNA) was used as a template. The first PCR reaction was diluted 1:20 and 1 μ l was used in a 50 μ l PCR reaction with SBE A and Ri primers and the products were cloned into pT7Blue. The cloned PCR products were screened for the presence or absence of the CSBE23 oligo by colony PCR.

A full length cDNA of cassava SBE II was isolated by PCR from leaf or root cDNA (RoRidT17 primed) using primers CSBE214 and CSBE218 from 2.5 μ l of cDNA in a 25 μ l reaction and 30 cycles of 94°C 45 sec, 55°C 25 sec, 72°C 2 min.

Complementation of E. coli mutant KV832

SBE II containing plasmids were transformed into the branching enzyme deficient mutant E.~coli~KV832 (Keil et al., 1987 Mol. Gen. Genet. 207, 294-301) and cells grown on solid PYG media (0.85 % KH₂PO₄, 1.1 % K₂HPO₄, 0.6 % yeast extract) containing 1.0 % glucose. To test for complementation, a loop of cells was scraped off and resuspended in 150 μ L water to which was added 15 μ L of Lugol's solution (2 g KI and 1 g I₂ per 300 ml water).

RNA isolation

RNA was isolated from cassava plants by the method of Logemann (1987 Anal. Biochem. 163, 21-26). Leaf RNA was isolated from 0.5 gm of in vitro grown plant tissue. The total yield was 300 μ g. Three month old roots (88 gm) were used for isolation of root RNA).

SBE II specific oligonucleotides

SBE A ATGGACAAGGATATGTATGA (Seq ID No. 1)

CSBE21 GGTTTCATGACTTCTGAGCA (Seq ID No. 2)

CSBE22	TGCTCAGAAGTCATGAAACC	(Seq ID No. 3)
CSBE23	TCCAGTCTCAATATACGTCG	(Seq ID No. 4)
CSBE24	AGGAGTAGATGGTCTGTCGA	(Seq ID No. 5)
CSBE25	TCATACATATCCTTGTCCAT	(Seq ID No. 6)
CSBE26	GGGTGACTTCAATGATGTAC	(Seq ID No. 7)
CSBE27	GGTGTACATCATTGAAGTCA	(Seq ID No. 8)
CSBE28	AATTACTGGCTCCGTACTAC	(Seq ID No. 9)
CSBE29	CATTCCAACGTGCGACTCAT	(Seq ID No. 10)
CSBE210	TACCGGTAATCTAGGTGTTG	(Seq ID No. 11)
CSBE211	GGACCTTGGTTTAGATCCAA	(Seq ID No. 12)
CSBE212	ATGAGTCGCACGTTGGAATG	(Seq ID No. 13)
CSBE213	CAACACCTAGATTACCGGTA	(Seq ID No. 14)
CSBE214	TTAGTTGCGTCAGTTCTCAC	(Seq ID No. 15)
CSBE215	AATATCTATCTCAGCCGGAG	(Seq ID No. 16)
CSBE216	ATCTTAGATAGTCTGCATCA	(Seq ID No. 17)
CSBE217	TGGTTGTTCCCTGGAATTAC	(Seq ID No. 18)
CSBE218	TGCAAGGACCGTGACATCAA	(Seq ID No. 19)

RESULTS

Cloning of a SBE II gene from cassava leaf

The strategy for cloning a full length cDNA of starch branching enzyme II of cassava is shown in Figure 1. A comparison of several SBE II (class A) SBE DNA sequences identified a 23 bp region which appears to be completely conserved among most genes (data not shown) and is positioned about one kilobase upstream from the 3' end of the gene. An oligonucleotide primer (designated SBE A) was made to this sequence and used to isolate a partial cDNA clone by 3' RACE PCR from first strand leaf cDNA as illustrated in Figure 1. An approximately 1100 bp band was amplified, cloned into pT7Blue vector and sequenced. This clone was designated pSJ94 and contained a 1120 bp insert starting with the SBE A oligo and ending with a polyA tail. There was a predicted open reading frame of 235 amino acids which was highly homologous (79% identical) to a potato SBE II also isolated by the inventors (data not shown) suggesting that this clone represented a class A (SBE II) gene.

To obtain the sequence of a full length clone nested primers were made complementary to the 5' end of this sequence and used in 5' RACE PCR to isolate clones from the 5' region of the gene. A total of three rounds of 5' RACE was needed to determine the sequence of the complete gene (i.e. one that has a predicted long ORF preceded by stop codons). It should be noted that during this cloning process several clones (# 23, 9, 16) were obtained that had small deletions and in one case (clone 23) there was also a small (120 bp) intron present. These occurrences are not uncommon and probably arise through errors in the PCR process and/or reverse transcription of incompletely processed RNA (heterogeneous nuclear RNA).

The overlapping cDNA fragments could be assembled into a contiguous 3 kb sequence (designated csbe2con.seq) which contained one long predicted ORF as shown in Figure 2. Several clones in the last round of 5' RACE were obtained which included sequence of the untranslated leader (UTL). All of these clones had an ORF (42 amino acids) 46 bp upstream and out of frame with that of the long ORF.

There is more than one SBE II gene in cassava

In order to determine if the assembled sequence represented that of a single gene, attempts were made to recover by PCR a full length SBE II gene using primers CSBE214 and CSBE23 at the 5' and 3' ends of the csbe2con sequence respectively. All attempts were unsuccessful using either leaf or root cDNA as template. The PCR was therefore repeated with either the 5'- or 3'- most primer and complementary primers along the length of the SBE II gene to determine the size of the largest fragment that could be amplified. With the CSBE214 primer, fragments could be amplified using primers 210, 28, 27 and 22 in order of increasing distance, the latter primer pair amplifying a 2.2 kb band. With the 3' primer CSBE23, only primer pairs with 21 and 26 gave amplification products, the latter being about 1200 bp. These results suggest that the original 3' RACE clone (pSJ94) is derived from a different SBE II gene than the rest of the 5' RACE clones even though the two largest PCR fragments (214+22 and 26+23) overlap by 750 bp and share several primer sites. It is likely that the sequence of the two genes starts to diverge around the CSBE22 primer site such that the 3' end of the corresponding gene does not contain the 23 primer and is not therefore able to amplify a cDNA when used with the 214 primer.

To confirm this, the sequence of the longest 5' PCR fragment (214+22) from two clones (#20 designated pSJ99, & #35) was determined and compared to the consensus sequence csbe2con as shown in Figure 3. The first 2000 bases are nearly identical (the single base changes might well be PCR errors), however the consensus sequence is significantly different after this. This region corresponds to the original 3' RACE fragment pSJ94 (SBE A + Ri adaptor) and provided evidence that there may be more than one SBE II gene in cassava.

The 3' end corresponding to pSJ99 was therefore cloned as follows: 3' RACE PCR was performed on leaf cDNA using the SBE A oligo as the gene specific primer so that all SBE II genes would be amplified. The cloned DNA fragments were then screened for the presence or absence of the CSBE23 primer by PCR. Two out of 15 clones were positive with the SBE A + Ri primer pair but negative with SBE A + CSBE23 primers. The sequence of these two clones (designated pSJ101, as shown in Figure 9) demonstrated that they were indeed from an SBE II gene and that they were different from pSJ94. However the overlapping region of pSJ101 (the 3' clone) and pSJ99 (the 5' clone) was identical suggesting that they were derived from the same gene.

To confirm this a primer (CSBE218) was made to a region in the 3' UTR (untranslated region) of pSJ101 and used in combination with CSBE214 primer to recover by PCR a full length cDNA from both leaf and root cDNA. These clones were sequenced and designated pSJ106 & pSJ107 respectively. The sequence and predicted ORF of pSJ107 is shown in Figure 4. The long ORF in plasmid pSJ106 was found to be interrupted by a stop codon (presumably introduced in the PCR process) approximately 1 kb from the 3' end of the gene, therefore another cDNA clone (designated pSJ116) was amplified in a separate reaction, cloned and sequenced. This clone had an intact ORF (data not shown). There were only a few differences in these two sequences (in the transit peptide aa 27-41: YRRTSSCLSFNFKEA to DRRTSSCLSFIFKKAA and L831 in pSJ107 to V in pSJ116 respectively).

An additional 740bp of sequence of the gene corresponding to the pSJ94 clone was isolated by 5' RACE using the primers CSBE216 and 217, and was designated pSJ125.

This sequence was combined with that of pSJ94 to form a consensus sequence "125 + 94", as shown in Figure 10. The sequence of this second gene is about 90% identical at the DNA and protein level to pSJ116, as shown in Figure 5 and 6, and is clearly a second form of SBE II in cassava. The 3' untranslated regions of the two genes are not related (data not shown).

It was also determined that the full length cassava SBE II genes (from both leaf and tuber) actually encode for active starch branching enzymes since the cloned genes were able to complement the glycogen branching enzyme deficient *E. coli* mutant KV832.

Main Findings

- 1) A full length cDNA clone of a starch branching enzyme II (SBE II) gene has been cloned from leaves and starch storing roots of cassava. This cDNA encodes a 836 amino acid protein (Mr 95 Kd) and is 86 % identical to pea SBE I over the central conserved domain, although the level of sequence identity over the entire coding region is lower than 86%.
- 2) There is more than one SBE II gene in cassava as a second partial SBE II cDNA was isolated which differs slightly in the protein coding region from the first gene and has no homology in the 3' untranslated region.
- 3) The isolated full length cDNA from both leaves and roots encodes an active SBE as it complements an *E. coli* mutant deficient in glycogen branching enzyme as assayed by iodine staining.

We have shown that there are SBE II (Class A) gene sequences present in the cassava genome by isolating cDNA fragments using 3' and 5' RACE. From these cDNA fragments a consensus sequence of over 3 kb could be compiled which contained one long open reading frame (Figure 2) which is highly homologous to other SBE II (class A) genes (data not shown). It is likely that the consensus sequence does not represent that of a single gene since attempts to PCR a full length gene using primers at the 5' and 3' ends of this sequence were not successful. In fact screening of a number of leaf derived 3'

RACE cDNAs showed that a second SBE II gene (clone designated pSJ101) was also expressed which is highly homologous within the coding region to the originally isolated cDNA (pSJ94) but has a different 3' UTR. A full length SBE II gene was isolated from leaves and roots by PCR using a new primer to the 3' end of this sequence and the original sequence at the 5' end of the consensus sequence. If the frequency of clones isolated by 3' RACE PCR reflects the abundance of the mRNA levels then this full length gene may be expressed at lower levels in the leaf than the pSJ94 clone (2 out of 15 were the former class, 13/15 the latter). It should be noted that each class is expressed in both leaves and roots as judged by PCR (data not shown). Sequence analysis of the predicted ORF of the leaf and root genes showed only a few differences (4 amino acid changes and one deletion) which could have arisen through PCR errors or, alternatively, there may be more than one nearly identical gene expressed in these tissues.

A comparison of all known SBE II protein sequences shows that the cassava SBE II gene is most closely related to the pea gene (Figure 8). The two proteins are 86.3% identical over a 686 amino acid range which extends from the triple proline "elbow" (Burton et al., 1995 Plant J. 7, 3-15) to the conserved VVYA sequence immediately preceding the Cterminal extensions (data not shown). All SBE II proteins are conserved over this range in that they are at least 80% similar to each other. Remarkably however, the sequence conservation between the pea, potato and cassava SBE II proteins also extends to the Nterminal transit peptide, especially the first 12 amino acids of the precursor protein and the region surrounding the mature terminus of the pea protein (AKFSRDS). Because the proteins are so similar around this region it can be predicted that the mature terminus of the cassava SBE II protein is likely to be GKSSHES. The precursor has a predicted molecular mass of 96 kD and the mature protein a predicted molecule mass of 91.3 kD. The cassava SBE II has a short acidic tail at the C-terminal although this is not as long or as acidic as that found in the pea or potato proteins. The significance of this acidic tail, if any, remains to be determined. One notable difference between the amino acid sequence of cassava SBE II and all other SBE II proteins is the presence of the sequence NSKH at around position 697 instead of the conserved sequence DAD/EY. Although this conserved region forms part of a predicted α -helix (number 8) of the catalytic $(\beta/\alpha)_8$ barrel domain (Burton et al 1995 cited previously), this difference does not abolish the SBE activity of the cassava protein as this gene can still complement the glycogen branching deletion mutant of *E. coli*. It may however affect the specificity of the protein. An interesting point is that the other cassava SBE II clone pSJ94 has the conserved sequence DADY.

One other point of interest concerning the sequence of the SBE II gene is the presence of an upstream ATG in the 5' UTR. This ATG could initiate a small peptide of 42 amino acids which would terminate downstream of the predicted initiating methionine codon of the SBE II precursor. If this does occur then the translation of the SBE II protein from this mRNA is likely to be inefficient as ribosomes normally initiate at the 5' most ATG in the mRNA. However the first ATG is in a poorer Kozak context than the SBE II initiator and it may be too close to the 5' end of the message to initiate efficiently (14 nucleotides) thus allowing initiation to occur at the correct ATG.

In conclusion we have shown that cassava does have SBE II gene sequences, that they are expressed in both leaves and tubers and that more than one gene exists.

Example 2 Cloning of a second full length cassava SBE II gene

Methods

Oligonucleotides

CSBE219	CTTTATCTATTAAAGACTTC	(Seq ID No. 20)
CSBE220	CAAAAAGTTTGTGACATGG	(Seq ID No. 21)
CSBE221	TCACTTTTTCCAATGCTAAT	(Seq ID No. 22)
CSBE222	TCTCATGCAATGGAACCGAC	(Seq ID No. 23)
CSBE223	CAGATGTCCTGACTCGGAAT	(Seq ID No. 24)
CSBE224	ATTCCGAGTCAGGACATCTG	(Seq ID No. 25)
CSBE225	CGCATTTCTCGCTATTGCTT	(Seq ID No. 26)
CSBE226	CACAGGCCCAAGTGAAGAAT	(Seq ID No. 27)

The 5' end of the gene corresponding to the 3'RACE clone pSJ94 was isolated in three

rounds of 5'RACE. Prior to performing the first round of 5' RACE, 5 μ g of total leaf RNA was reverse transcribed in a 20 μ l reaction using conditions as decribed by the manufacturer (Superscript enzyme, BRL) and 10 pmol of the SBE II gene specific primer CSBE23. Primers were then removed and the cDNA tailed with dATP as described above. The first round of 5'RACE used primers CSBE216 and Ro. This PCR reaction was diluted 1:20 and used as a template for a second round of amplification using primers CSBE217 and Ri. The gene specific primers were designed so that they would preferentially hybridise to the SBE II sequence in pSJ94. Amplified products appeared as a smear of approximately 600-1200 bp when subjected to electrophoresis on a 1% TAE agarose gel.

This smear was excised and DNA purified using a Qiaquick column (Qiagen) before ligation to the pT7Blue vector. Several clones were sequenced and clone #7 was designated pSJ125. New primers (CSBE219 and 220) were designed to hybridise to the 5' end of pSJ125 and a second round of 5'RACE was performed using the same CSBE23 primed library. Two fragments of 600 and 800 bp were cloned and sequenced (clones 13,17). Primers CSBE221 and 222 were designed to hybridise to the 5' sequence of the longest clone (#13) and a third round of 5' RACE was performed on a new library (5 µg total leaf RNA reverse transcribed with Superscript using CSBE220 as primer and then dATP tailed with TdT from Boehringer Mannheim). Fragments of approximately 500 bp were amplified, cloned and sequenced. Clone #13, was designated pSJ143. The process is illustrated schematically in Figure 12.

To isolate a full length gene as a contiguous sequence, a new primer (CSBE225) was designed to hybridise to the 5' end of clone pSJ143 and used with one of the primers (CSBE226 or 23) in the 3' end of clone pSJ94, in a PCR reaction using RoRidT17 primed leaf cDNA as template. Use of primer CSBE226 resulted in production of Clone #2 (designated pSJ144), and use of primer CSBE23 resulted in production of Clones #10 and 13 (designated pSJ145 and pSJ146 respectively). Only pSJ146 was sequenced fully.

Results

Isolation of a second full length cassava SBE II gene

A full length clone for a second SBE II gene was isolated by extending the sequence of pSJ94 in three rounds of 5' RACE as illustrated schematically in Figure 12. In each round of 5' RACE, primers were designed that would preferentially hybridise to the new sequence rather than to the gene represented by pSJ116. In the final round of 5' RACE, three clones were obtained that had the initiating methione codon, and none of these had upstream ATGs. The overlapping cDNA fragments (sequences of the 5'RACE clones pSJ143, 13, pSJ125 and the 3'RACE clone pSJ94) could be assembled into a consensus sequence of approximately 3 kb which was designated csbe2-2.seq. This sequence contained one long ORF with a predicted size of 848 aa (M_r 97 kDa). The full length gene was then isolated as a contiguous sequence by PCR amplification from RoRidT17 primed leaf cDNA using primers at the 5' (CSBE225) and 3' (CSBE23 or CSBE226) ends of the RACE clones. One clone, designated pSJ146, was sequenced and the restriction map is shown along with the predicted amino acid sequence in Figure 13.

Sequence homologies between SBE II genes

The two cassava genes (pSJ116 and pSJ146) share 88.8% identity at the DNA level over the entire coding region (data not shown). The homology extends about 50 bases outside of this region but beyond this the untranslated regions show no similarity (data not shown). At the protein level the two genes show 86% identity over the entire ORF (data not shown). The two genes are more closely related to each other than to any other SBE II. Between species, the pea SBE I shows the most homology to the cassava SBE II genes.

Example 3

Construction of plant transformation vectors and transformation of cassava with antisense starch branching enzyme genes.

This example describes in detail how a portion of the SBE II gene isolated from cassava may be introduced into cassava plants to create transgenic plants with altered properties.

An 1100 bp *Hind* III - *Sac* I fragment of cassava SBE II (from plasmid pSJ94) was cloned into the *Hind* III - *Sac* I sites of the plant transformation vector pSJ64 (Figure 11). This placed the SBE II gene in an antisense orientation between the 2X 35S CaMV promoter

and the nopaline synthase polyadenylation signal. pSJ64 is a derivative of the binary vector pGPTV-HYG (Becker et al., 1992 Plant Molecular Biology 20: 1195-1197) modified by inclusion of an approximately 750 bp fragment of pJIT60 (Guerineau et al 1992 Plant Mol. Biol. 18, 815-818) containing the duplicated cauliflower mosaic virus (CaMV) 35S promoter (Cabb-JI strain, equivalent to nucleotides 7040 to 7376 duplicated upstream of 7040 to 7433, as described by Frank et al., 1980 Cell 21, 285-294) to replace the GUS coding sequence. A similar construct was made with the cassava SBE II sequence from plasmid pSJ101.

These plasmids are then introduced into Agrobacterium tumefaciens LBA4404 by a direct DNA uptake method (An et al., Binary vectors, In: Plant Molecular Biology Manual (ed Galvin and Schilperoort) AD 1988 pp 1-19) and can be used to transform cassava somatic embryos by selecting on hygromycin as described by Li et al. (1996, Nature Biotechnology 14, 736-740).

SEQUENCE LISTING

(1) GENERAL INFORMATION:	
 (i) APPLICANT: (A) NAME: National Starch and Chemical Investment	
(ii) TITLE OF INVENTION: Improvements in or Relating to Starch Content of Plants	
(iii) NUMBER OF SEQUENCES: 31	
<pre>(iv) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)</pre>	
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AAT Asn	GTA Val 60	ATG Met	GTC Val	ACT Thr	GCT Ala	TCT Ser 65	AAA Lys	AGA Arg	GTC Val	CTT Leu	CCT Pro 70	GAT Asp	GGT Gly	CGG Arg	ATT Ile		242
GAA G1u 75	TGC Cys	TAT Tyr	TCT Ser	TCT Ser	TCA Ser 80	ACA Thr	GAT Asp	CAA Gln	TTG Leu	GAA G1u 85	GCC Ala	CCT Pro	GGC Gly	ACA Thr	GTT Val 90		290
	GAA Glu																338
GAT Asp	AAG Lys	ATT Ile	GTT Val 110	GAA Glu	GAT Asp	GAA Glu	GTA Val	AAT Asn 115	AAA Lys	GAA Glu	TCT Ser	GTT Val	CCA Pro 120	ATG Met	CGG Arg		386
	ACA Thr																434
	CCC Pro 140																482
GGC Gly 155	TTT Phe	CGT Arg	CAA Gln	CAC His	CTA Leu 160	GAT Asp	TAC Tyr	CGG Arg	TAT Tyr	TCA Ser 165	CAG Gln	TAC Tyr	AAA Lys	AGA Arg	CTC Leu 170		530
	GAA Glu																578
	TAT Tyr															·	626
	GAG Glu																674
	AAC Asn 220																722

GTC Val 235	TGG Trp	GAG Glu	ATC Ile	TTT Phe	TTG Leu 240	CCG Pro	AAT Asn	AAT Asn	GCA Ala	GAT Asp 245	GGT Gly	TCA Ser	CCA Pro	CCA Pro	ATT Ile 250	770
CCC Pro	CAT His	GGT Gly	TCT Ser	CGA Arg 255	GTA Val	AAG Lys	ATA Ile	CGC Arg	ATG Met 260	GAT Asp	ACT Thr	CCA Pro	TCT Ser	GGC Gly 265	AAC Asn	· <u>81</u> 8
AAA Lys	GAT Asp	TCT Ser	ATT Ile 270	CCT Pro	GCT Ala	TGG Trp	ATC- Ile	AAG Lys 275	TTC Phe	TCA Ser	GTT Val	CAA Gln	GCA Ala 280	CCA Pro	GGT Gly	866
GAA Glu	CTC Leu	CCA Pro 285	TAT Tyr	AAT Asn	GGC Gly	ATA Ile	TAC Tyr 290	TAT Tyr	GAT Asp	CCT Pro	CCC Pro	GAG G1u 295	GAG Glu	GAG Glu	AAG Lys	914
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TAT Tyr 315	GAG G1u	TCG Ser	CAC His	GTT Val	GGA Gly 320	ATG Met	AGT Ser	AGT Ser	ACG Thr	GAG Glu 325	CCA Pro	GTA Val	ATT Ile	AAC Asn	ACA Thr 330	1010
TAT Tyr	GCC Ala	AAC Asn	TTT Phe	AGA Arg 335	GAT Asp	GAT Asp	GTG Val	CTT Leu	CCT Pro 340	CGC Arg	ATC Ile	AAA Lys	AAG Lys	CTT Leu 345	GGC Gly	1058
TAC Tyr	AAT Asn	GCT Ala	GTT Val 350	CAG Gln	CTC Leu	ATG Met	GCT Ala	ATT Ile 355	CAA Gln	GAG Glu	CAT His	TCA Ser	TAT Tyr 360	TAT Tyr	GCT Ala	1106
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GGA Gly	ACT Thr 380	CCT Pro	GAT Asp	GAT Asp	TTA Leu	AAG Lys 385	TCT Ser	CTA Leu	ATA Ile	GAT Asp	AAA Lys 390	GCT Ala	CAC His	GAG Glu	TTA Leu	1202
GGT Gly 395	CTT Leu	CTT Leu	GTT Val	CTC Leu	ATG Met 400	GAT Asp	ATT Ile	GTT Val	CAT His	AGC Ser 405	CAT His	GCA Ala	TCA Ser	ACT Thr	AAT Asn 410	1250
											GAT Asp					1298
											GAC Asp					1346
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ACT Thr 475	TCA Ser	ATG Met	ATG Met	TAC Tyr	ACC Thr 480	CAT His	CAT His	GGA Gly	TTG Leu	CAG G1n 485	GTA Val	GAT Asp	TTT Phe	ACC Thr	GGC Gly 490	- mar	1490
AAC Asn	TAC Tyr	AAT Asn	GAA Glu	TAC Tyr 495	TTT Phe	GGA Gly	TAT Tyr	GCA Ala	ACT Thr 500	GAT Asp	GTA Val	GAT Asp	GCT Ala	GTG Val 505	GTT Val		1538
TAT Tyr	TTG Leu	ATG Met	CTG Leu 510	TTG Leu	AAT Asn	GAT Asp	ATG Met	ATT Ile 515	CAT His	GGT Gly	CTC Leu	TTC Phe	CCA Pro 520	GAG Glu	GCT Ala		1586
GTC Val	ACC Thr	ATT Ile 525	GGT Gly	GAA Glu	GAT Asp	GTT Val	AGT Ser 530	GGA Gly	ATG Met	CCA Pro	ACA Thr	GTT Val 535	TGC Cys	ATT Ile	CCG Pro		1634
GTT Val	GAA G1u 540	GAT Asp	GGT Gly	GGT Gly	GTT Val	GGC Gly 545	TTT Phe	GAT Asp	TAT Tyr	CGT Arg	CT C Leu 550	CAC His	ATG Met	GCT Ala	GTT Val		1682
GCT A1a 555	GAT Asp	AAA Lys	TGG Trp	GTT Val	GAG G1u 560	ATT Ile	ATT Ile	CAG Gln	AAG Lys	AGA Arg 565	GAT Asp	GAA Glu	GAT Asp	TGG Trp	AAA Lys 570		1730
ATG Met	GGT Gly	GAC Asp	ATT Ile	GTA Val 575	CAT His	ATG Met	CTG Leu	ACC Thr	AAC Asn 580	AGG Arg	CGG Arg	TGG Trp	TTG Leu	GAA Glu 585	AAG Lys		1778
TGT Cys	GTT Val	TCT Ser	TAT Tyr 590	GCT Ala	GAA Glu	AGT Ser	CAT His	GAC Asp 595	CAG Gln	GCC Ala	CTT Leu	GTT Val	GGT Gly 600	GAC Asp	AAA Lys		1826
ACT Thr	ATT Ile	GCA Ala 605	TTT Phe	TGG Trp	CTG Leu	ATG Met	GAC Asp 610	AAG Lys	GAT Asp	ATG Met	TAT Tyr	GAC Asp 615	TTC Phe	ATG Met	GCT Ala		1874
CTT Leu	GAC Asp 620	AGA Arg	CCA Pro	TCT Ser	ACT Thr	CCT Pro 625	CTC Leu	ATA Ile	GAT Asp	CGT Arg	GGA Gly 630	GTA Val	GCA Ala	TTG Leu	CAC His		1922
AAA Lys 635	ATG Met	ATC Ile	AGG Arg	CTT Leu	ATT Ile 640	ACC Thr	ATG Met	GGA Gly	TTA Leu	GGC Gly 645	GGA Gly	GAA Glu	GGA Gly	TAT Tyr	TTG Leu 650		1970
AAT Asn	TTT Phe	ATG Met	GGA Gly	AAT Asn 655	GAA Glu	TTT Phe	GGA Gly	CAC His	CCC Pro 660	GAG Glu	TGG Trp	ATT Ile	GAT Asp	TTT Phe 665	CCA Pro		2018
AGA Arg	GGT Gly	GAT Asp	CTA Leu 670	CAT His	CTT Leu	CCC Pro	AGT Ser	GGT Gly 675	AAA Lys	TTT Phe	GTT Val	CCT Pro	GGG Gly 680	AAC Asn	AAT Asn		2066

TAC Tyr	AGT Ser	TAT Tyr 685	GAT Asp	AAA Lys	TGC Cys	CGG Arg	CGT Arg 690	AGG Arg	TTT Phe	GAT Asp	CTA Leu	GGC Gly 695	AAT Asn	TCA Ser	AAG Lys	2114
CAT His	CTG Leu 700	AGA Arg	TAT Tyr	CAT His	GGA Gly	ATG Met 705	CAA Gln	GAG Glu	TTT Phe	GAT Asp	CAA Gln 710	GCA Ala	ATT	CAG Gln	CAT His	2162
CTT Leu 715	GAA Glu	GAA Glu	GCC Ala	TAT Tyr	GGT Gly 720	TTC Phe	ATG Met	ACT Thr	TCT Ser	GAG Glu 725	CAC His	CAA Gln	TAC Tyr	ATA Ile	TCA Ser 730	2210
CGG Arg	AAG Lys	GAT Asp	GAA Glu	AGG Arg 735	GAT Asp	CGG Arg	ATC Ile	ATT Ile	GTC Val 740	TTC Phe	GAG Glu	AGG Arg	GGA Gly	AAC Asn 745	ETC Leu	2258
GTT Val	TTT Phe	GTA Val	TTC Phe 750	AAT Asn	TTT Phe	CAT His	TGG Trp	ACT Thr 755	AGC Ser	AGC Ser	TAT Tyr	TCG Ser	GAT Asp 760	TAC Tyr	CGA Arg	2306
GTT Val	GGC Gly	TGC Cys 765	TTA Leu	AAG Lys	CCA Pro	GGA Gly	AAG Lys 770	TAC Tyr	AAG Lys	ATA Ile	GTC Val	TTG Leu 775	GAT Asp	TCA Ser	GAT Asp	2354
GAT Asp	CCT Pro 780	TTG Leu	TTT Phe	GGA Gly	GGC Gly	TTT Phe 785	GGC Gly	AGG Arg	CTT Leu	AGT Ser	CAT His 790	GAT Asp	GCA Ala	GAG Glu	CAC His	2402
TTC Phe 795	AGC Ser	TTT Phe	GAA Glu	GGG Gly	TGG Trp 800	TAC Tyr	GAT Asp	AAC Asn	CGG Arg	CCT Pro 805	CGA Arg	TCC Ser	TTC Phe	ATG Met	GTG Val 810	2450
TAC Tyr	ACA Thr	CCA Pro	TGT Cys	AGA Arg 815	ACA Thr	GCA Ala	GTG Val	GTC Val	TAT Tyr 820	GCT Al-a	TTA Leu	GTG Val	GAG Glu	GAT Asp 825	GAA Glu	2498
GTG Val	GAG G1u	AAT Asn	GAA Glu 830	TTG Leu	GAA Glu	CCT Pro	GTC Val	GCC Ala 835	GGT Gly	TAA *	GATA	ATAT(CTT A	VACA/	ACAGGT	2551
TCT	TCTGAAGCAG GAATGCCATT ATTGATCTTC CTATGTT												2588			

(2) INFORMATION FOR SEQ ID NO: 29:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 837 amino acids
 - (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

Met Gly His Tyr Thr Ile Ser Gly Ile Arg Phe Pro Cys Ala Pro Leu $1 \hspace{1cm} 5 \hspace{1cm} . \hspace{1cm} 10 \hspace{1cm} - \hspace{1cm} 15$

Cys Lys Ser Gln Ser Thr Gly Phe His Gly Tyr Arg Arg Thr Ser Ser Cys Leu Ser Phe Asn Phe Lys Glu Ala Phe Ser Arg Arg Val Phe Ser Gly Lys Ser Ser His Glu Ser Asp Ser Ser Asn Val Met Val Thr Ala 50 55 60 Ser Lys Arg Val Leu Pro Asp Gly Arg Ile Glu Cys Tyr Ser Ser Ser 65 70 75 80 Thr Asp Gln Leu Glu Ala Pro Gly Thr Val Ser Glu Glu Ser Gln Val Leu Thr Asp Val Glu Ser Leu Ile Met Asp Asp Lys Ile Val Glu Asp 105 100 Glu Val Asn Lys Glu Ser Val Pro Met Arg Glu Thr Val Ser Ile Arg Lys Ile Gly Ser Lys Pro Arg Ser Ile Pro Pro Pro Gly Arg Gly Gln 130 140 Arg Ile Tyr Asp Ile Asp Pro Ser Leu Thr Gly Phe Arg Gln His Leu Asp Tyr Arg Tyr Ser Gln Tyr Lys Arg Leu Arg Glu Glu Ile Asp Lys
165 170 175 Tyr Glu Gly Ser Leu Asp Ala Phe Ser Arg Gly Tyr Glu Lys Phe Gly 180 Phe Ser Arg Ser Glu Thr Gly Ile Thr Tyr Arg Glu Trp Ala Pro Gly 200 Ala Thr Trp Ala Ala Leu Ile Gly Asp Phe Asn Asn Trp Asn Pro Asn 210 215 220 210 Ala Asp Val Met Thr Gln Asn Glu Cys Gly Val Trp Glu Ile Phe Leu 230 Pro Asn Asn Ala Asp Gly Ser Pro Pro Ile Pro His Gly Ser Arg Val 245 250 255 Lys Ile Arg Met Asp Thr Pro Ser Gly Asn Lys Asp Ser Ile Pro Ala 260 265 270 260 Trp Ile Lys Phe Ser Val Gln Ala Pro Gly Glu Leu Pro Tyr Asn Gly 280 Ile Tyr Tyr Asp Pro Pro Glu Glu Glu Lys Tyr Val Phe Lys Asn Pro 290 295 300 Gln Pro Lys Arg Pro Lys Ser Leu Arg Ile Tyr Glu Ser His Val Gly 310 305

Met Ser Ser Thr Glu Pro Val Ile Asn Thr Tyr Ala Asn Phe Arg Asp 330 Asp Val Leu Pro Arg Ile Lys Lys Leu Gly Tyr Asn Ala Val Gln Leu Met Ala Ile Gln Glu His Ser Tyr Tyr Ala Ser Phe Gly Tyr His Val Thr Asn Phe Tyr Ala Ala Ser Ser Arg Phe Gly Thr Pro Asp Asp Leu Lys Ser Leu Ile Asp Lys Ala His Glu Leu Gly Leu Leu Val Leu Met Asp-Ile Val His Ser His Ala Ser Thr Asn Thr Leu Asp Gly Leu Asn 410 Met Phe Asp Gly Thr Asp Gly His Tyr Phe His Ser Gly Pro Arg Gly His His Trp Met Trp Asp Ser Arg Leu Phe Asn Tyr Gly Ser Trp Glu 435 440 445 Val Leu Arg Phe Leu Leu Ser Asn Ala Arg Trp Trp Leu Asp Glu Tyr Lys Phe Asp Gly Phe Arg Phe Asp Gly Val Thr Ser Met Met Tyr Thr 47Ŏ His His Gly Leu Gln Val Asp Phe Thr Gly Asn Tyr Asn Glu Tyr Phe Gly Tyr Ala Thr Asp Val Asp Ala Val Val Tyr Leu Met Leu Leu Asn Asp Met Ile His Gly Leu Phe Pro Glu Ala Val Thr Ile Gly Glu Asp 515 Val Ser Gly Met Pro Thr Val Cys Ile Pro Val Glu Asp Gly Gly Val Gly Phe Asp Tyr Arg Leu His Met Ala Val Ala Asp Lys Trp Val Glu 545 560 IIe Ile Gln Lys Arg Asp Glu Asp Trp Lys Met Gly Asp Ile Val His 565 570 575 Met Leu Thr Asn Arg Arg Trp Leu Glu Lys Cys Val Ser Tyr Ala Glu Ser His Asp Gln Ala Leu Val Gly Asp Lys Thr Ile Ala Phe Trp Leu Met Asp Lys Asp Met Tyr Asp Phe Met Ala Leu Asp Arg Pro Ser Thr 615

Pro Leu Ile Asp Arg Gly Val Ala Leu His Lys Met Ile Arg Leu Ile 625 630 635 640

Thr Met Gly Leu Gly Gly Glu Gly Tyr Leu Asn Phe Met Gly Asn Glu 645 650

Phe Gly His Pro Glu Trp Ile Asp Phe Pro Arg Gly Asp Leu His Leu 660 665 670

Pro Ser Gly Lys Phe Val Pro Gly Asn Asn Tyr Ser Tyr Asp Lys Cys 675 680 685

Arg Arg Arg Phe Asp Leu Gly Asn Ser Lys His Leu Arg Tyr His Gly 690 700

Met Gln Glu Phe Asp Gln Ala Ile Gln His Leu Glu Glu Ala Tyr Gly 705 710 715 720

Phe Met Thr Ser Glu His Gln Tyr Ile Ser Arg Lys Asp Glu Arg Asp 725 730 735

Arg Ile Ile Val Phe Glu Arg Gly Asn Leu Val Phe Val Phe Asn Phe 740 745 750

His Trp Thr Ser Ser Tyr Ser Asp Tyr Arg Val Gly Cys Leu Lys Pro
755 760 _ 765

Gly Lys Tyr Lys Ile Val Leu Asp Ser Asp Asp Pro Leu Phe Gly Gly 770 780

Phe Gly Arg Leu Ser His Asp Ala Glu His Phe Ser Phe Glu Gly Trp 785 790 795 800

Tyr Asp Asn Arg Pro Arg Ser Phe Met Val Tyr Thr Pro Cys Arg Thr 805 810 815

Ala Val Val Tyr Ala Leu Val Glu Asp Glu Val Glu Asp Glu Leu Glu 820 825 830

Pro Val Ala Gly * 835

(2) INFORMATION FOR SEQ ID NO: 30:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2805 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION:131..2677
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

AGTGAATTCG AGCTCGGTAC CCGGGGATCC GATTCGCATT TCTCGCTATT GCTTTCCGTT 60												
TATTTCCATA TATAAAATAT CAAATCTAAT CACTTGCGCC ATTTCTATCT CTCTCCAAAC 1												
TCTCACCGAA ATG Met		Thr Val Ser G	GGC ATA CGT T Gly Ile Arg P 345		S							
GCA CCT TCA CTC Ala,Pro Ser Leu												
AGG ACC TCT TCT Arg Thr Ser Ser 870	Gly Leu Ser		Lys Lys Glu									
CGG AAG ATC TTT Arg Lys Ile Phe 885												
TTA ACT GTC TCT Leu Thr Val Ser 900												
GAT GGC TCT TCT Asp Gly Ser Ser 915				Gly Thr V								
TTG GAG GAA TCC Leu Glu Glu Ser												
GAT GAT AAG AAT Asp Asp Lys Asr 950	ı Val Glu Glu		Lys Lys Glu									
TTG CAT GAG ACA Leu His Glu Thr 965												
ATT CCT CCA CCT The Pro Pro Pro 980												
TTG GCA GGT TTC Leu Ala Gly Phe 995				Gln Tyr L								
AGG CTG CGT GAG Arg Leu Arg Glu			Gly Gly Leu									
TCT CGT GGA TT Ser Arg Gly Phe 103	e Glu Lys Phe		Arg Ser Glu									

30	
ACT TAT AGG GAA TGG GCA CCT GGA GCT ACG TGG GCT GCA CTT ATT GGA Thr Tyr Arg Glu Trp Ala Pro Gly Ala Thr Trp Ala Ala Leu Ile Gly 1045 1050 1055	793
GAT TTC AAC AAT TGG AAT CCT AAT GCA GAT GTC ATG ACT CGG AAT GAG Asp Phe Asn Asn Trp Asn Pro Asn Ala Asp Val Met Thr Arg Asn Glu 1060 1070	841
TTT GGT GTC TGG GAG ATT TTT TTG CCA AAT AAC GCA GAT GGT TCA CCA Phe Gly Val Trp Glu Ile Phe Leu Pro Asn Ala Asp Gly Ser Pro 1075 1080 1085 1090	889
CCA ATT CCT CAT GGT TCT CGA GTA AAG ATA CGC ATG GAT ACT CCA TCT Pro Ile Pro His Gly Ser Arg Val Lys Ile Arg Met Asp Thr Pro Ser 1095	937
GGC ATC AAA GAT TCA ATT CCT GCT TGG ATC AAG TTC TCA GTT CAG GCA Gly Ile Lys Asp Ser Ile Pro Ala Trp Ile Lys Phe Ser Val Gln Ala - 1110 1115 1120	985
CCT GGT GAA ATC CCA TAC AAT GCC ATA TAC TAT GAT CCA CCA AAG GAG Pro Gly Glu Ile Pro Tyr Asn Ala Ile Tyr Tyr Asp Pro Pro Lys Glu 1125 1130 1135	1033
GAG AAG TAT GTG TTC AAA CAT CCT CAG CCA AAG AGA CCA AAA TCA CTT Glu Lys Tyr Val Phe Lys His Pro Gln Pro Lys Arg Pro Lys Ser Leu 1140 1145 1150	1081
AGG ATT TAT GAA TCT CAT GTT GGG ATG AGT AGT ATG GAG CCA ATA ATT Arg Ile Tyr Glu Ser His Val Gly Met Ser Ser Met Glu Pro Ile Ile 1155 1160 1165 1170	1129
AAC ACA TAT GCC AAC TIT AGA GAT GAT ATG CTT CCT CGC ATC AAA AAG Asn Thr Tyr Ala Asn Phe Arg Asp Asp Met Leu Pro Arg Ile Lys Lys 1175 1180 1185	1177
CTT GGC TAC AAT GCT GTT CAG ATC ATG GCT ATT CAA GAG CAT TCC TAT Leu Gly Tyr Asn Ala Val Gln Ile Met Ala Ile Gln Glu His Ser Tyr 1190 1195 1200	1225
TAT GCT AGT TTT GGG TAC CAT GTC ACA AAC TTT TTT GCA CCT AGC AGC Tyr Ala Ser Phe Gly Tyr His Val Thr Asn Phe Phe Ala Pro Ser Ser 1205 1210 1215	1273
CGA TTT GGA ACT CCT GAT GAT TTG AAG TCT TTA ATA GAT AAA GCT CAT Arg Phe Gly Thr Pro Asp Asp Leu Lys Ser Leu Ile Asp Lys Ala His 1220 1230	1321
GAG TTA GGG CTG CTT GTT CTC ATG GAT ATT GTT CAT AGC CAT GCG TCA Glu Leu Gly Leu Leu Val Leu Met Asp Ile Val His Ser His Ala Ser 1235 1240 1245 1250	1369
AAT AAT ACG TTG GAT GGG CTG AAC ATG TTT GAT GGT ACG GAT AGT CAC Asn Asn Thr Leu Asp Gly Leu Asn Met Phe Asp Gly Thr Asp Ser His 1255	1417

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TAC Tyr	TTC Phe	CAC His	TCC Ser 127	ыу	TCA Ser	CGG Arg	GGT Gly	CAT His 127	His	TGG Trp	TTG Leu	TGG Trp	GAC Asp 128	Ser	CGC Arg	1465
CTT Leu	TTC Phe	AAC Asn 128	Tyr	GGA Gly	AGC Ser	TGG Trp	GAG Glu 129	Val	CTA Leu	AGA Arg	TTT Phe	CTT Leu 129	Leu	TCA Ser	AAT Asn	1513
GCA Ala	AGA Arg 1300	1rp	TGG Trp	TTG Leu	GAA Glu	GAG Glu 130	Tyr	AGG Arg	TTT Phe	GAT Asp	GGT Gly 1310	Phe	AGA Arg	TTT Phe	GAT Asp	1561
GGG Gly 1319	Val	ACT Thr	TCC Ser	ATG Met	ATG Met 1320	lyr	ACT Thr	CCC Pro	CAT His	GGG Gly 1325	Leu	CAG Gln	GTA Val	GCT Ala	TTT Phe 1330	1609
ACT Thr	GGC Gly	AAC Asn	TAC Tyr	AAT Asn 133	Glu	TAC Tyr	TTT Phe	GGA Gly	TAT Tyr 1340	Ala	ACT Thr	GAT Asp	GTA Val	GAT Asp 134	Ala	1657
GTG Val	ATT Ile	TAT Tyr	TTG Leu 1350	Met	CTT Leu	GTG Val	AAT Asn	GAT Asp 1359	Met	ATT Ile	CAC His	GGT Gly	CTT Leu 1360	Phe	CCT Pro	1705
GAG G1u	GCT Ala	GTT Val 1365	ACC Thr	ATT Ile	GGT Gly	GAA Glu	GAT Asp 1370	Val	AGC Ser	GGA Gly	AAG Lys	CCA Pro 1375	Thr	TTT Phe	TGC Cys	1753
ATT Ile	CCA Pro 1380	Val	GAA Glu	GAT Asp	GGT Gly	GGT Gly 1385	Val	GGA Gly	TTT Phe	GAT Asp	TAC Tyr 1390	Arg	CTC Leu	CAC His	ATG Met	1801
GCC Ala 1395	He	GCC Ala	GAT Asp	AAA Lys	TGG Trp 1400	He	GAG G1u	ATT Ile	CTT Leu	AAG Lys 1405	Lys	AGA Arg	GAT Asp	GAG Glu	GAC Asp 1410	1849
TGG Trp	AAA Lys	ATG Met	GGT Gly	GAC Asp 1415	He	GTG Val	CAT His	ACA Thr	CTC Leu 1420	Thr	AAC Asn	AGA Arg	AGG Arg	TGG Trp 1425	Leu	1897
GAA G1u	AAA Lys	TGT Cys	GTT Val 1430	Ala	TAT Tyr	GCT Ala	GAA G1u	AGT Ser 1435	His	GAC Asp	CAA Gln	GCT Ala	CTT Leu 1440	۷a٦	GG ^T Gly	1945
GAC Asp	AAA Lys	ACT Thr 1445	ATT Ile	GCA Ala	TTT Phe	TGG Trp	CTG Leu 1450	Met	GAC Asp	AAG Lys	GAC Asp	ATG Met 1455	Tyr	GAC Asp	TTC Phe	1993
Met	GCT Ala 1460	Arg	GAC Asp	AGA Arg	CCA Pro	TCT Ser 1465	Thr	CCT Pro	CTT Leu	Ile	GAT Asp 1470	Arg	GGA Gly	ATA Ile	GCA Ala	2041
TTG Leu 1475	HIS	AAA Lys	ATG Met	ATC Ile	AGG Arg 1480	Leu	ATT Ile	ACC Thr	ATG Met	GGC Gly 1485	Leu	GGC Gly	GGA Gly	GAA Glu	GGA Gly 1490	2089

									40							-
TAT T Tyr L	TG A. .eu A.	AT T sn P	ne	ATG Met 1495	GIY	AAT Asn	GAA Glu	TTT Phe	GGA Gly 150	His	CCT Pro	GAG Glu	TGG Trp	ATT Ile 150	Asp	2137
TTT C Phe P	CCA A(Pro Ai	rg u	GG 11 y 510	ASP	CGA Arg	CAT His	CTG Leu	CCC Pro 151	Asn	GGT Gly	AAA Lys	GTA Val	ATT Ile 152	Pro	GGG Gly	2185
AAC A Asn (A	Sn H	AC A is S 525	GT er	TAT Tyr	GAT Asp	AAA Lys	TGC Cys 1530	Arg	CGT Arg	AGA Arg	TTT Phe	GAT Asp 1539	Leu	GGT Gly	GAT Asp	2233
GCA GA Ala As 15	AC TA sp Ty 540	AT C yr L	TA . eu .	AGA Arg	TAT Tyr	CAT His 1545	Gly	ATG Met	CAA Gln	GAG Glu	TTT Phe 1550	Asp	CAG Gln	GCA Ala	ATG Met	2281
CAA C/ Gln H 1555	AT CT is Le	∏ G eu G	AA (]u (GAA Glu	GCC Ala 1560	Tyr	GGT Gly	TTC Phe	ATG Met	ACT Thr 1565	Ser	GAG G1u	CAC His	CAG Gln	TAT Tyr 1570	2329
ATA TO	CA CG er Ar	GG A	ys /	GAT Asp 1575	Glu	GGA Gly	GAT Asp	CGG Arg	ATC Ile 1580	Ile	GTC Val	TTT Phe	GAG Glu	AGG Arg 1585	Gly	2377
AAC CT Asn Le	TT GT eu Va	al Pi	TT (he 1 590	GTA Val	TTC Phe	AAC Asn	TTT Phe	CAT His 1595	Trp	ACT Thr	AAC Asn	AGC Ser	TAT Tyr 1600	Ser	GAT Asp	2425
TAC CO Tyr Ar	rg va	T G(11 G 305	GC ⁻ ly (TGC Cys	TTC Phe	AAG Lys	TCA Ser 1610	Gly	AAG Lys	TAC Tyr	AAG Lys	ATT Ile 1615	Val	TTG Leu	GAC Asp	2473
TCG GA Ser As	AT GA sp As 620	AT G(sp G	GC T	TTG _eu	Phe	GGA Gly 1625	Gly	TTC Phe	AAC Asn	AGG Arg	CTT Leu 1630	Ser	CAT His	GAT Asp	GCC Ala	2521
GAG CA Glu Hi 1635	AC TT is Ph	C A(ie Th	CC T	he .	GAC Asp 1640	Gly	TGG Trp	TAT Tyr	GAT Asp	AAC Asn 1 6 45	Arg	CCT Pro	CGG Arg	TCC Ser	TTC Phe 1650	2569
ATG GT Met Va	TA TA al Ty	T G(r Al	Ia F	CCA Pro : 1655	TCT Ser	AGG Arg	ACA Thr	Ala	GTG Val 1660	Val	TAT Tyr	GCT Ala	TTA Leu	GTA Val 1665	Glu	2617
GAT GA Asp G1	AA GA lu Gl	u As	AT (sn (570	SAA (Slu /	GCA Ala	GAG G1u	Asn	GAA Glu 1675	Val	GAA Glu	AGT Ser	Glu	GTG Val 1680	Lys	CCA Pro	2665
GCC TC Ala Se	CC GG er Gl 16	у *	A G	GATA(GATA	TTT	AGTA	AGAG	G AT	CCCC	TAAA	GCA	GGAA	TGG		2717
TTAACC	CTGTG	CAT	ГСТС	CAT	T GA	ACGA	CGTA	TAT	TGAG	ACT	GGAA	ATCC	AT A	TGAC	TAGTA	2777
GATCCT	ГСТАG	AGT	CG/	ACCT	G CA	GGCA	TG							-	-	-2805

(2) INFORMATION FOR SEQ ID NO: 31:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 849 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

Met Val Tyr Tyr Thr Val Ser Gly Ile Arg Phe Pro Cys Ala Pro Ser 1 5 10 15

Leu Tyr Lys Ser Gln Leu Thr Ser Phe His Gly Gly Arg Arg Thr Ser 25 30

Ser Gly Leu Ser Phe Leu Leu Lys Lys Glu Leu Phe Pro Arg Lys Ile 35 40 45

Phe Ala Gly Lys Ser Ser Tyr Glu Ser Asp Ser Ser Asn Leu Thr Val 50 55 60

Ser Ala Ser Glu Lys Val Leu Val Pro Asp Asp Gln Ile Asp Gly Ser 65 70 75 80

Ser Ser Ser Thr Tyr Gln Leu Glu Thr Thr Gly Thr Val Leu Glu Glu 85 90 95

Ser Gln Val Leu Gly Asp Ala Glu Ser Leu Val Met Glu Asp Asp Lys 100 110

Asn Val Glu Glu Asp Glu Val Lys Lys Glu Ser Val Pro Leu His Glu 115 120 125

Thr Ile Ser Ile Gly Lys Ser Glu Ser Lys Pro Arg Ser Ile Pro Pro 130 135 140

Pro Gly Ser Gly Gln Arg Ile Tyr Asp Ile Asp Pro Ser Leu Ala Gly 145 150 155 160

Phe Arg Gln His Leu Asp Tyr Arg Tyr Ser Gln Tyr Lys Arg Leu Arg 165 170 175

Glu Glu Ile Asp Lys Tyr Glu Gly Gly Leu Asp Ala Phe Ser Arg Gly 180 185 190

Phe Glu Lys Phe Gly Phe Leu Arg Ser Glu Thr Gly Ile Thr Tyr Arg 200 205

Glu Trp Ala Pro Gly Ala Thr Trp Ala Ala Leu Ile Gly Asp Phe Asn 210 215 220

Asn Trp Asn Pro Asn Ala Asp Val Met Thr Arg Asn Glu Phe Gly Val 225 235 240

Trp Glu Ile Phe Leu Pro Asn Asn Ala Asp Gly Ser Pro Pro Ile Pro His Gly Ser Arg Val Lys Ile Arg Met Asp Thr Pro Ser Gly Ile Lys 265 270 Asp Ser Ile Pro Ala Trp Ile Lys Phe Ser Val Gln Ala Pro Gly Glu 275 280 285 Ile Pro Tyr Asn Ala Ile Tyr Tyr Asp Pro Pro Lys Glu Glu Lys Tyr 290 295 300 Val Phe Lys His Pro Gln Pro Lys Arg Pro Lys Ser Leu Arg Ile Tyr 305 310 315 320 Glu Ser His Val Gly Met Ser Ser Met Glu Pro Ile Ile Asn Thr Tyr 325 330 335 Ala Asn Phe Arg Asp Asp Met Leu Pro Arg Ile Lys Lys Leu Gly Tyr Asn Ala Val Gln Ile Met Ala Ile Gln Glu His Ser Tyr Tyr Ala Ser 355 360 365 Phe Gly Tyr His Val Thr Asn Phe Phe Ala Pro Ser Ser Arg Phe Gly 375 Thr Pro Asp Asp Leu Lys Ser Leu Ile Asp Lys Ala His Glu Leu Gly Leu Leu Val Leu Met Asp Ile Val His Ser His Ala Ser Asn Asn Thr 405 Leu Asp Gly Leu Asn Met Phe Asp Gly Thr Asp Ser His Tyr Phe His 420 425 430 Ser Gly Ser Arg Gly His His Trp Leu Trp Asp Ser Arg Leu Phe Asn 440 Tyr Gly Ser Trp Glu Val Leu Arg Phe Leu Leu Ser Asn Ala Arg Trp Trp Leu Glu Glu Tyr Arg Phe Asp Gly Phe Arg Phe Asp Gly Val Thr 475 470 480 Ser Met Met Tyr Thr Pro His Gly Leu Gln Val Ala Phe Thr Gly Asn Tyr Asn Glu Tyr Phe Gly Tyr Ala Thr Asp Val Asp Ala Val Ile Tyr 500 505 Leu Met Leu Val Asn Asp Met Ile His Gly Leu Phe Pro Glu Ala Val 520 Thr Ile Gly Glu Asp Val Ser Gly Lys Pro Thr Phe Cys Ile Pro Val 530 540



Glu Asp Gly Gly Val Gly Phe Asp Tyr Arg Leu His Met Ala Ile Ala Asp Lys Trp Ile Glu Ile Leu Lys Lys Arg Asp Glu Asp Trp Lys Met 565 570 575 Gly Asp Ile Val His Thr Leu Thr Asn Arg Arg Trp Leu Glu Lys Cys Val Ala Tyr Ala Glu Ser His Asp Gln Ala Leu Val Gly Asp Lys Thr 595 600 605 Ile Ala Phe Trp Leu Met Asp Lys Asp Met Tyr Asp Phe Met Ala Arg 610 620 Asp Arg Pro Ser Thr Pro Leu Ile Asp Arg Gly Ile Ala Leu His Lys 625 630 635 640 Met Ile Arg Leu Ile Thr Met Gly Leu Gly Gly Glu Gly Tyr Leu Asn 645 650 655 Phe Met Gly Asn Glu Phe Gly His Pro Glu Trp Ile Asp Phe Pro Arg Gly Asp Arg His Leu Pro Asn Gly Lys Val Ile Pro Gly Asn Asn His 675 680 685 Ser Tyr Asp Lys Cys Arg Arg Phe Asp Leu Gly Asp Ala Asp Tyr **69**5 Leu Arg Tyr His Gly Met Gln Glu Phe Asp Gln Ala Met Gln His Leu 705 710 715 720 Glu Glu Ala Tyr Gly Phe Met Thr Ser Glu His Gln Tyr Ile Ser Arg 725 730 735 Lys Asp Glu Gly Asp Arg Ile Ile Val Phe Glu Arg Gly Asn Leu Val 740 745 750 Phe Val Phe Asn Phe His Trp Thr Asn Ser Tyr Ser Asp Tyr Arg Val. 765 Gly Cys Phe Lys Ser Gly Lys Tyr Lys Ile Val Leu Asp Ser Asp Asp 770 780 --Gly Leu Phe Gly Gly Phe Asn Arg Leu Ser His Asp Ala Glu His Phe 785 790 795 800 Thr Phe Asp Gly Trp Tyr Asp Asn Arg Pro Arg Ser Phe Met Val Tyr 805 810 815 Ala Pro Ser Arg Thr Ala Val Val Tyr Ala Leu Val Glu Asp Glu Glu 820 825 830 Asn Glu Ala Glu Asn Glu Val Glu Ser Glu Val Lys Pro Ala Ser Gly

SUBSTITUTE SHEET (RULE 26)